```
chain nodes :
7 8 9 11 29
ring nodes :
1 2 3 4 5 6 10 12 13 14 15 17 18 19 20 21 22 23 24 25 26 27
28
chain bonds :
3-7 7-8 8-9 8-11 9-10 14-17 28-29
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6 10-12 10-15 12-13 13-14 14-15 17-18 17-22
18-19 19-20 20-21 21-22 23-28 23-24 24-25 25-26 26-27 27-28
exact/norm bonds :
3-7 7-8 8-9 8-11 9-10 10-12 10-15 12-13 13-14 14-15 28-29
exact bonds :
14-17
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6 17-18 17-22 .18-19 19-20 20-21 21-22 23-28
23-24 24-25 25-26 26-27 27-28
isolated ring systems :
containing 1 : 10 : 17 : 23 :
Match level :
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:Atom
11:CLASS 12:Atom 13:Atom 14:Atom 15:Atom 17:Atom 18:Atom 19:Atom 20:Atom
```

21:CLASS 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:Atom 28:Atom 29:CLASS

L2 STRUCTURE UPLOADED

=> d L2 HAS NO ANSWERS L2 STR *** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation.

30:CLASS

L4 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2006 ACS on STN
RN 478365-32-1 REGISTRY
ED Entered STN: 08 Jan 2003
Urea, N-[2-pheny]-4-thiazoly])-N'-[6-[4-(1-pyrrolidinylmethyl)phenoxy]-2pyridinyl]- [9C1) (CA INDEX NAME)
S 3D CONCORD
MF C26 H25 N5 02 S
CA
LC STN Files: CA, CAPLUS, TOXCENTER, USPATZ, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

Page 8 ELHILO

=> d ibib abs hitstr tot 'IBIB' IS NOT A VALID FORMAT FOR FILE 'REGISTRY' 'ABS' IS NOT A VALID FORMAT FOR FILE 'REGISTRY' 'HITSTR' IS NOT A VALID FORMAT FOR FILE 'REGISTRY' The following are valid formats: Substance information can be displayed by requesting individual fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number) REG - RN - Index Name, MF, and structure - no RN SAM FIDE - All substance data, except sequence data - FIDE, but only 50 names SQIDE - IDE, plus sequence data SQIDE3 - Same as SQIDE, but 3-letter amino acid codes are used - Protein sequence data, includes RN SQD - Same as SQD, but 3-letter amino acid codes are used SQD3 SON - Protein sequence name information, includes RN CALC - Table of calculated properties EPROP - Table of experimental properties PROP - EPROP and CALC Any CA File format may be combined with any substance format to obtain CA references citing the substance. The substance formats must be cited first. The CA File predefined formats are: ABS -- Abstract APPS -- Application and Priority Information BIB -- CA Accession Number, plus Bibliographic Data CAN -- CA Accession Number CBIB -- CA Accession Number, plus Bibliographic Data (compressed) IND -- Index Data IPC -- International Patent Classification PATS -- PI, SO STD -- BIB, IPC, and NCL IABS -- ABS, indented, with text labels IBIB -- BIB, indented, with text labels ISTD -- STD format, indented OBIB ----- AN, plus Bibliographic Data (original) OIBIB ----- OBIB, indented with text labels SBIB ----- BIB, no citations SIBIB ----- IBIB, no citations it is available.

The ALL format gives FIDE BIB ABS IND RE, plus sequence data when

The MAX format is the same as ALL.

The IALL format is the same as ALL with BIB ABS and IND indented, with text labels.

For additional information, please consult the following help

messages:

HELP DFIELDS -- To see a complete list of individual display fields. HELP FORMATS -- To see detailed descriptions of the predefined formats. ENTER DISPLAY FORMAT (IDE):ide

```
L4 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2006 ACS on STN
RN 478365-32-1 REGISTRY
ED Entered STN: 09 Jan 2003
Urea, N-(2-phenyl-4-thiazoly1)-N'-[6-[4-(1-pyrrolidinylmethyl)phenoxy]-2-
pyridinyl)- (9Cl) (CA INDEX NAME)
S 3D CONCORD
MF C26 H2S N5 02 S
CA
LC STN Files: CA, CAPLUS, TOXCENTER, USPATZ, USPATFULL
```

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)